```
(FILE 'MEDLINE, CANCERLIT, BIOTECHDS, EMBASE, CAPLUS, BIOSIS' ENTERED AT
     11:29:35 ON 17 MAR 2003)
                DEL HIS
          39240 S MICROPARTICLE OR MICROSPHERE
L1
         306151 S EMULSI?
L2
L3
         106208 S EVAPORAT?
           587 S L3 AND L2 AND L1
L4
           2425 S PEG AND DEXTRAN
L5
L6
              3 S L5 AND L4
L7
             1 DUP REM L6 (2 DUPLICATES REMOVED)
         740447 S DISPERS? OR DISCONTINUOUS
L8
             95 S L8 AND L4
L9
             72 S L9 AND PHASE
L10
             37 DUP REM L10 (35 DUPLICATES REMOVED)
L11
L12
        3101431 S DNA OR NUCLEIC OR PLASMID
             15 S L12 AND L4
L13
             9 DUP REM L13 (6 DUPLICATES REMOVED)
L14
             11 S L5 AND L3
L15
             5 DUP REM L15 (6 DUPLICATES REMOVED)
L16
             1 S PEG AND L2 AND L1 AND L12 AND AQUEOUS
L17
            491 S PEG AND L1
L18
            92 S L18 AND L8
L19
L20
             5 S L19 AND L12
L21
             5 DUP REM L20 (0 DUPLICATES REMOVED)
```

=>

L7 ANSWER 1 OF 1 MEDLINE

AN 97402448 MEDLINE

DN 97402448 PubMed ID: 9259512

Biodegradable polymeric microparticles for drug delivery and vaccine formulation: the surface attachment of hydrophilic species using the concept of poly(ethylene glycol) anchoring segments.

DUPLICATE 1

- AU Coombes A G; Tasker S; Lindblad M; Holmgren J; Hoste K; Toncheva V; Schacht E; Davies M C; Illum L; Davis S S
- CS Department of Pharmaceutical Sciences, University of Nottingham, University Park, UK.
- SO BIOMATERIALS, (1997 Sep) 18 (17) 1153-61. Journal code: 8100316. ISSN: 0142-9612.
- CY ENGLAND: United Kingdom
- DT Journal; Article; (JOURNAL ARTICLE)
- LA English
- FS Priority Journals
- EM 199712
- ED Entered STN: 19980109
 Last Updated on STN: 19980109
 Entered Medline: 19971202
- Poly(ethylene glycol)-dextran (PEG-DEX) conjugates AB have been used as a combined stabilizer and surface modifier to produce resorbable poly(DL-lactide-co-glycolide) (PLG) microparticles by an emulsification/solvent evaporation technique. The use of PEG or dextran polymers alone was incapable of producing microparticles. Particle size measurements revealed smaller mean particle sizes (480 nm) and improved polydispersity when using a 1.2% PEG substituted conjugate relative to a 9% substituted material (680 nm). PLG microparticles modified by post-adsorbed PEG-DEX conjugates flocculated in 0.01 M salt solutions, whereas PLG microparticles prepared using PEG-DEX as a surfactant were stable in at least 0.5 M NaCl solutions. Surface modification of PLG microparticles was confirmed by zeta potential measurements and surface analysis using X-ray photoelectron spectroscopy. The presence of surface exposed dextran was confirmed by an immunological detection method using a dextran -specific antiserum in an enzyme-linked immunosorbent assay. The findings support a model in which the PEG component of the PEG -DEX conjugate provides an anchor to the microparticle surface while the dextran component extends from the particle surface to contribute a steric stabilization function. This approach offers opportunities for attaching hydrophilic species such as targeting moieties to biodegradable microparticles to improve the interaction of drug carriers and vaccines with specific tissue sites.

L11 ANSWER 28 OF 37 EMBASE COPYRIGHT 2003 ELSEVIER SCI. B.V.

AN 96339585 EMBASE

DN 1996339585

TI Development of furosemide loaded polymethylmethacrylate microspheres.

AU Sa B.; Ghosh A.

CS Dept. of Pharmaceutical Technology, Jadavpur University, Calcutta - 700 032, India

SO Indian Drugs, (1996) 33/10 (521-524). ISSN: 0019-462X CODEN: INDRBA

CY India

DT Journal; Article

FS 037 Drug Literature Index

LA English

SL English

AB Furosemide-loaded polymethylmethacrylate microparticles were prepared by emulsion-solvent evaporation technique using aqueous phase as dispersion medium to minimize the use of hazardous organic solvents. Among the different emulsion stabilizers, methylcellulose and hydroxy propylcellulose, in a concentration range of 0.0025 to 0.1% w/v were found suitable for the formation of discrete and spherical microparticles. The spherical shape of the microparticles was not influenced by variation in concentration of either methylcellulose or hydroxy propylcellulose. Actual drug content in the microparticles was almost equal to theoretical drug load and was not influenced significantly by change in either methylcellulose or hydroxy-propylcellulose concentrations.

L14 ANSWER 8 OF 9 MEDLINE

MEDLINE

1999093470 ANDN

99093470 PubMed ID: 9874713

- PLGA microspheres containing plasmid DNA: preservation TIof supercoiled DNA via cryopreparation and carbohydrate stabilization.
- Ando S; Putnam D; Pack D W; Langer R ΑU
- Massachusetts Institute of Technology, E25-342, 45 Carleton Street, CS Cambridge, Massachusetts 02139, USA.

DUPLICATE 4

- SO JOURNAL OF PHARMACEUTICAL SCIENCES, (1999 Jan) 88 (1) 126-30. Journal code: 2985195R. ISSN: 0022-3549.
- CY United States
- Journal; Article; (JOURNAL ARTICLE) DT
- LΑ English
- FS Priority Journals
- 199903 EM
- ED Entered STN: 19990402

Last Updated on STN: 19990402

Entered Medline: 19990322

AΒ Biodegradable microspheres containing plasmid DNA have potential uses as mediators of transfection in cells, particularly phagocytic cells such as macrophages. However, the hydrophilic nature and the structural instability of supercoiled DNA preclude its facile encapsulation in polymer matrixes such as poly(d, l-lactic-co-glycolic acid) (PLGA) by traditional methods. We initially studied the microencapsulation of plasmid DNA using the established water-in-oil-in-water double-emulsion solventevaporation method and found that (1) the encapsulation efficiency was low (about 20%), (2) the microencapsulation procedure nicked (degraded) the supercoiled DNA, and (3) lyophilization of the microsphere also nicked the DNA. We have therefore designed a new microsphere preparation method (called cryopreparation) to specifically address these concerns. Using the cryopreparation method, the aqueous phase of the primary emulsion containing the plasmid DNA is frozen and then subjected to homogenization. Because there is no shear stress inside a solid, we hypothesized that freezing the aqueous phase of the primary emulsion would help to preserve the supercoiled plasmid DNA during formation of the secondary emulsion. We also hypothesized that the formation of crystals from buffers within the primary emulsion was a causative factor for nicking during freezing or lyophilization, and that disruption of the crystal formation by the addition of saccharides into the primary emulsion would improve the supercoiled-DNA content of the spheres. Our results support the two hypotheses. Not only was the supercoiled-DNA content increased from 39% to over 85%, but the encapsulation efficiency was also elevated from 23% to over 85%.

ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
ANSWER 5 OF 9 CAPLUS COPYRIGHT 2003 ACS
     2001:380725 CAPLUS
AN
DN
     135:2561
TI
     Continuous-flow method for preparing microparticles
IN
     Hedley, Mary Lynne; Hsu, Yung-Yueh; Tyo, Michael
PA
     Zycos, Inc., USA
     PCT Int. Appl., 47 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LΑ
     English
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
                                           -----
     ______
                      ----
                            _____
                                       WO 2000-US31770 20001117
PΙ
     WO 2001036583
                     A1 20010525
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
             YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                      A1 20020814 EP 2000-978814 20001117
     EP 1230338
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
PRAI US 1999-443654
                     A1 19991119
     WO 2000-US31770
                      W
                            20001117
     The invention is based on the discovery of a method for scalable,
AΒ
     continuous flow prodn. of a nucleic acid-contg.
     microparticle that maintains the structural integrity of the
     assocd. nucleic acid and results in a microparticle
     having a purity suitable for introduction into an animal (e.g., human)
     host. Microparticles prepd. according to the continuous flow processes
     described herein can be used for delivery of a nucleic acid for
     gene therapy, antisense therapy, vaccination, treatment of autoimmune
     disease, and either specific or non-specific modulation of an immune
     response (e.g., via cytokine regulation). The microparticles can addnl.
     be used to deliver nucleic acid encoding a protein or peptide
     useful in any type of therapy.
```

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE.CNT 3

```
L21 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2003 ACS
     1995:546956 CAPLUS
AN
DN
     122:274119
TI
     Hydrophobic polymeric pharmaceutical microparticles
     Andrianov, Alexander K.; Langer, Robert S.
IN
PA
     Virus Research Institute, USA; Massachusetts Institute of Technology
SO
     PCT Int. Appl., 33 pp.
     CODEN: PIXXD2
DT
     Patent
LΑ
     English
FAN.CNT 1
     PATENT NO.
                  KIND DATE
                                          APPLICATION NO. DATE
                     ----
                           -----
     WO 9508320
                     A1
                            19950330
                                         WO 1994-US10692 19940921
ΡI
         W: AM, AU, BB, BG, BR, BY, CA, CN, CZ, FI, GE, HU, JP, KG, KP, KR,
             KZ, LK, LT, LU, LV, MD, MG, MN, MW, NO, NZ, PL, RO, RU, SI, SK,
            TJ, TT, UA, UZ, VN
         RW: KE, MW, SD, SZ, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU,
            MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN,
             TD, TG
     US 5500161
                            19960319
                                          US 1993-124816
                                                            19930921
     CA 2172040
                           19950330
                      AA
                                          CA 1994-2172040 19940921
    AU 9478001
                           19950410
                                          AU 1994-78001
                      A1
                                                           19940921
     EP 720471
                                          EP 1994-928640
                      Α1
                           19960710
                                                           19940921
     EP 720471
                      В1
                           20010418
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE
                                          AT 1994-928640
     AT 200616
                      \mathbf{E}
                           20010515
                                                          19940921
     ES 2159305
                      Т3
                           20011001
                                          ES 1994-928640
                                                           19940921
PRAI US 1993-124816
                      Α
                           19930921
    WO 1994-US10692
                     W
                           19940921
AΒ
    A method for the prepn. of microparticles, and the product thereof, that
     include dispersing a substantially water insol. non-ionic or
     ionic polymer in an aq. soln. in which the substance to be delivered is
     also dissolved, dispersed or suspended, and then coagulating the
     polymer together with the substance by impact forces to form a
    microparticle. In an alternative embodiment, the
    microparticle is formed by coagulation of an aq. polymeric
     dispersion through the use of electrolytes, pH changes, org.
     solvents in low concns. (the minimal amt. necessary to break up the
     dispersion), or temp. changes to form polymer matrixes
     encapsulating biol. materials. Thus 60 mg of fluorescein-labeled bovine
     serum albumin was dissolved in 3 mL of 30% aq. soln. dispersion
     of Eudragite NE 30D and then spraying the ag. dispersion in a
     flask contg. 200 mL of deionized water using Turbotack air-atomizing
    nozzle. The flow rate of the polymeric dispersion was
     150.mu.L/min, the air pressure was 25 psi, and the distance between the
    nozzle and surface of water was 30 cm. The resulting microparticles were
     spherical with an av. diam. of 1-10 .mu.m and encapsulation efficiency of
     65%.
```

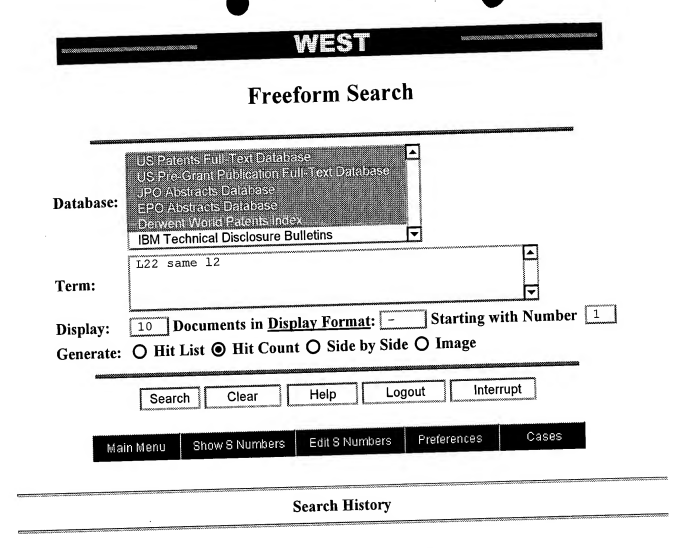
malonate) microparticles with encapsulated paclitaxel were prepd.

```
L21 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2003 ACS
AN
     2002:654974 CAPLUS
DN
     137:190745
TI
     Emulsion-based processes for making microparticles
IN
     Gibson, John W.; Holl, Richard J.; Tipton, Arthur J.
PΑ
     Southern Biosystems, Inc., USA
SO
     U.S., 16 pp., Cont.-in-part of U.S. 6,291,013.
     CODEN: USXXAM
DT
     Patent
LА
     English
FAN.CNT 2
     PATENT NO.
                  KIND DATE
                                            APPLICATION NO. DATE
     ----
                      ____
                                            US 6440493 B1 20020827
                                            US 2000-726108 20001129
     US 2002142093
                      A1 20021003
     US 6291013 B1 20010918
WO 2000066087 A2 20001109
                                            US 1999-303842
                                                              19990503
                                          WO 2000-US11781 20000502
         W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,
             CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI,
             SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ,
             BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
             DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
             CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRAI US 1999-303842
                      A2 19990503
     WO 2000-US11781
                       A2
                           20000502
     Processes for making microparticles, preferably contg. an active agent,
AΒ
     are provided. In a preferred embodiment, the process involves prepg. (1)
     a dispersed phase contg. an agent in a soln. of polymer and a
     first solvent; (2) a continuous phase contg. a surfactant, and a second
     solvent that is totally or partially immiscible with the first solvent;
     and (3) an extn. phase that is a nonsolvent for the polymer, a solvent for
     the continuous phase components, and a solvent for the first solvent,
    wherein the first solvent has soly. in the extn. phase of between about
    0.1% and 25% by wt. Then, the dispersed phase and the
    continuous phase are mixed to form an emulsion, and the emulsion is then
    briefly mixed with a suitable quantity of extn. phase to induce skin
    formation at the interface of the dispersed and continuous
    phases. Remaining solvent is removed by an evapn. process step. The
    emulsification and solvent removal steps are preferably conducted in a
    continuous process. The brief extn. step prior to evapn. minimizes the
    loss of active agent from the microparticles, and reduces the required
    vol. of extn. phase as compared to other extn.-based processes. Alternate
    emulsification methods and solvent removal methods, such as incremental
    extn., cryogenic extn., or membrane sepn., also are provided, and can be
```

used in various combinations to make microparticles. Fluorescein-loaded

THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD

microspheres were prepd. using polylactide.



DATE: Monday, March 17, 2003 Printable Copy Create Case

aunt	Sat	Na

Set Name	Query	Hit Count	
side by side		_	result set
	PB,JPAB,EPAB,DWPI; PLUR=YES; OP=ADJ		
<u>L23</u>	L22 same l2	7	<u>L23</u>
<u>L22</u>	L21 same 117	112	<u>L22</u>
<u>L21</u>	l6 with peg	986	<u>L21</u>
<u>L20</u>	L18 same l1	12	<u>L20</u>
<u>L19</u>	L18 with 14	2	<u>L19</u>
<u>L18</u>	117 with 12	354	<u>L18</u>
<u>L17</u>	emulsif\$	176389	<u>L17</u>
<u>L16</u>	12 with 13	1	<u>L16</u>
<u>L15</u>	18 and 15	1	<u>L15</u>
<u>L14</u>	18 same 14	0	<u>L14</u>
<u>L13</u>	18 same 15	0	<u>L13</u>
<u>L12</u>	18 with 15	0	<u>L12</u>
<u>L11</u>	18 same 13	0	<u>L11</u>
<u>L10</u>	18 with 13	0	<u>L10</u>
<u>L9</u>	18 with 13L8	0	<u>L9</u>
<u>L8</u>	11 with 12	35	<u>L8</u>
<u>L7</u>	11 with 12 with 13 with 15	0	<u>L7</u>
<u>L6</u>	dextran	39963	<u>L6</u>
<u>L5</u>	evaporat?	61571	<u>L5</u>
<u>L4</u>	peg	70026	<u>L4</u>
<u>L3</u>	emulsif?	7308	<u>L3</u>
<u>L2</u>	microparticle	15154	<u>L2</u>
<u>L1</u>	dispersed phase	10292	<u>==</u> <u>L1</u>
	-		

END OF SEARCH HISTORY